

Brussels, 18.11.2021 C(2021) 8520 (final)

## **COMMISSION IMPLEMENTING DECISION**

of 18.11.2021

on the annual renewal of the conditional marketing authorisation for the orphan medicinal product for human use "Tecartus - autologous anti-CD19-transduced CD3+cells", granted by Decision C(2020) 9284(final)

(Text with EEA relevance)

(ONLY THE DUTCH TEXT IS AUTHENTIC)

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### THE EUROPEAN COMMISSION,

Having regard to the Treaty on the Functioning of the European Union,

Having regard to Regulation (EC) No 726/2004 of the European Parliament and of the Council of 31 March 2004 laying down Community procedures for the authorisation and supervision of medicinal products for human and veterinary use and establishing a European Medicines Agency<sup>1</sup>, and in particular Article 10(2) and 14-a thereof,

Having regard to Commission Regulation (EC) No 507/2006 on the conditional marketing authorisation for medicinal products for human use falling within the scope of Regulation (EC) No 726/2004 of the European Parliament and of the Council<sup>2</sup>,

Having regard to the application submitted by Kite Pharma EU B.V., on 11 June 2021, under Article 6(2) of Regulation (EC) No 507/2006 with a view to the annual renewal of the conditional marketing authorisation for the medicinal product "Tecartus - autologous anti-CD19-transduced CD3+ cells",

Having regard to the opinion of the European Medicines Agency, formulated on 16 September 2021 by the Committee for Medicinal Products for Human Use,

#### Whereas:

- (1) The medicinal product "Tecartus autologous anti-CD19-transduced CD3+ cells", entered in the Union Register of Medicinal Products under the number EU/1/20/1492 and authorised by Commission Decision C(2020) 9284(final) of 14 December 2020, remains in compliance with the requirements of Article 14-a of Regulation (EC) No 726/2004 of the European Parliament and of the Council, and Regulation (EC) No 507/2006,
- (2) The conditional marketing authorisation granted by Decision C(2020) 9284(final) should therefore be renewed.
- (3) The Union Register of Medicinal Products should be updated.
- (4) The measures provided for in this Decision are in accordance with the opinion of the Standing Committee on Medicinal Products for Human Use,

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OJ L 136, 30.4.2004, p. 1.

OJ L 92, 30.3.2006, p. 6.

## HAS ADOPTED THIS DECISION:

## Article 1

The conditional marketing authorisation granted by Decision C(2020) 9284(final) of 14 December 2020 is renewed without amendments.

### Article 2

The period of validity of the renewed authorisation shall be one year from 15 December 2021.

### Article 3

This Decision is addressed to Kite Pharma EU B.V., Tufsteen 1, 2132 NT Hoofddorp, Nederland.

Done at Brussels, 18.11.2021

For the Commission Sandra GALLINA Director-General